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On some forgotten species of *Exidia* and *Myxarium* (Auriculariales, Basidiomycota)

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Abstract

Exidia cartilaginea and *E. villosa* are typified and shown to be conspecific with the North American species *E. candida*. Based on morphological differences, *E. cartilaginea* is retained as a variety of *E. candida*. Current concept of *Myxarium nucleatum* covers four species in Nordic countries: *M. cinnamomescens*, *M. hyalinum*, *M. nucleatum* s.str. and *M. populinum*. These species are redescribed and their nomenclature is discussed. *Exidia subrepanda* is placed to the synonyms of *E. saccharina*, and *Heterochaete europaea* to *E. thuretiana*.

Introduction

The so-called jelly fungi (heterobasidiomycetes) are so far a deficiently studied group, especially at the generic and species levels. This paper deals with two genera, *Exidia* and *Myxarium*, currently considered members of the *Auriculariales* (Agaricomycetes). Despite certain morphological similarity, these genera are not closely related, and they belong to different lineages within the order (Weiss & Oberwinkler 2001, Hibbett et al. 2014). In Nordic countries, Neuhoﬀ's monograph (Neuhoﬀ 1936) was the last systematic attempt to revise species of *Exidia* s. l. (incl. *Myxarium*). Therefore, taxonomic revision based on modern methods and newly collected material is highly desirable. Here we investigate *Exidia*-like species with pale-coloured, orbicular or cerebriform basidiocarps, occurring on deciduous trees in North Europe.

First microscopy-based studies of this group were undertaken by Karsten. He regarded all pale-coloured exidioid fungi as belonging to one species, *Tremella albida* Huds. or *Exidia albida* (Huds.) Bref., and sequentially introduced f. *betulae* (Karsten 1866), subsp. *tuberculata* (Karsten 1889), var. *populina* and subsp. *subrepanda* (Karsten 1891). Oudemans (1919, 1920) accidentally raised the two latter taxa to the species rank but they remained completely ignored by all subsequent mycologists. Neuhoﬀ (1935) described two new species, *Exidia cartilaginea* S. Lundell & Neuhoﬀ and *E. villosa* Neuhoﬀ; a year later, he accepted two other species, *E. albida* and *E. gemmata* (Lév.) Bourdot & Maire, and reported all of them from Sweden (Neuhoﬀ 1936). Unfortunately, type specimens of *E. cartilaginea* and *E. villosa* kept in Königsberg were destroyed at the end of the Second World War (Torkelsen 1968, Jahn 1971). While the first species was well-known and widely treated from North Europe (Raitviir 1967, Torkelsen 1972, Hansen & Knudsen 1997), *E. villosa* stayed almost totally forgotten. Donk (1958) replaced *E. gemmata* with an older name *Myxarium nucleatum* Wallr. and then synonymized the latter one with *Myxarium hyalinum* (Pers.) Donk (based on *Tremella hyalina* Pers.) (Donk 1966). This solution aroused a long-term controversy (see, for example, Reid 1970 and Roberts 1998). Although recent DNA-based studies

confirmed an independent generic status of *Myxarium* (Weiss & Oberwinkler 2001, Wells et al. 2004), identity of both *M. hyalinum* and *M. nucleatum* has not been properly re-established. Competing or obscure names and uncertain species concepts are the source of insufficient knowledge of *Exidia* s.l. in North Europe.

In the present paper, we select type specimens for nine taxa. In total, five species and one variety are re-described and illustrated, and two new combinations in genera *Exidia* and *Myxarium* are proposed.

Material and methods

Type specimens and collections from herbaria H, O, LE, UPS, C, BPI, FH, TAAM, CWU were studied. Herbarium acronyms are given according to Thiers (2017). Morphological study follows Miettinen et al. (2012). The following abbreviations are used in morphological descriptions: L – mean spore length, W – mean spore width, Q – length/width ratio, n – number of measurements per specimens.

DNA extraction and amplification. For DNA extraction, small fragments of dried basidiocarps were used. In total, 27 specimens were selected for molecular sampling (Table 1). The procedure of DNA extraction completely corresponded to the manufacturing protocol of the NucleoSpin Plant II Kit (Macherey-Nagel GmbH & Co. KG). The following primers were used for both amplification and sequencing: ITS1F-ITS4 (<http://www.biology.duke.edu/fungi/mycolab/primers.htm>) for the nuclear ribosomal internal transcribed spacer (ITS) region; primers JS1 (Landvik 1996) and LR5 (<http://www.biology.duke.edu/fungi/mycolab/primers.htm>) for the large ribosomal subunit (nrLSU) region; primers EF1-983F and EF1-1567R for approximately 500bp of translation elongation factor 1- α (*tef1*) region (Rehner & Buckley 2005). PCR products were purified applying the GeneJET Gel Extraction Kit (Thermo Scientific, Thermo Fisher Scientific Inc., MA, USA). Sequencing was

performed with an ABI model 3130 Genetic Analyzer (Applied Biosystems, CA, USA). Raw data were edited and assembled in MEGA 6 (Tamura et al. 2013).

Phylogenetic analysis. For this study, 25 ITS, 26 nrLSU and 13 *tef1* sequences were generated (Table 1). The additional 11 ITS and 14 nrLSU sequences including the outgroups were retrieved from GenBank (www.ncbi.nlm.nih.gov/genbank/). Sequences were aligned with the MAFFT version 7 web tool (<http://mafft.cbrc.jp/alignment/server/>) using the Q-INS-i option for ITS and nrLSU. The final ITS+nrLSU alignment contained 1557 characters (including gaps). The final *tef1* alignment contained 523 characters (including gaps).

Phylogenetic reconstructions for data sets were performed with maximum likelihood (ML) and Bayesian (BA) analyses. Before the analyses, the best-fit substitution models for the alignment were estimated for each dataset based on Akaike Information Criterion (AIC) using FindModel web server (<http://www.hiv.lanl.gov/content/sequence/findmodel/findmodel.html>).

For all datasets, GTR model was selected. ML analysis was run in the PhyML server, v. 3.0 (<http://www.atgc-montpellier.fr/phyml/>), with one hundred rapid bootstrap replicates. BA was performed with MrBayes 3.1 software (Ronquist & Huelsenbeck 2003), for two independent runs, each with 5 million generations with sampling every 100 generations and four chains.

Newly generated sequences have been deposited in GenBank with corresponding accession numbers (Table 1).

Results

ML and BA analyses produced nearly the same topologies for the combined nrLSU – ITS dataset (Fig. 1). Specimens corresponding to the current concepts of *E. candida*, *E. cartilaginea* and *E. villosa* possess only negligible differences in their ITS and nrLSU sequences, and thus they should be considered conspecific. *tef1* dataset for these taxa show more variation which is, however, not congruent to any morphological, ecological or geographic traits. We designate this clade as *E.*

candida because this species name have priority versus the two others (see below). *Exidia candida* is rather distantly related to the similarly looking *E. thuretiana*, as well as to other *Exidia* species. A DNA-based revision of *Exidia* and *Exidiopsis* s.l. is desirable to properly define genus limits. All European collections of *Exidia*-like fungi with pedunculate basidia were so far considered as belonging to one species, *Myxarium nucleatum*. However, our data show that four species are hidden under this name (Fig. 1). They are clearly different in their DNA sequences, as well as in morphology and, to some degree, host preferences and geographic distribution. In the phylogenetic tree, these species are intermixed with totally effused (corticoid), non-jelly *Myxarium* species (*M. grilletii*, *M. subhyalinum*). This is, possibly, an indication that exidioid basidiocarps appeared independently in several lineages within this genus.

Species are re-described and commented in alphabetical order.

Exidia candida Lloyd, Mycol. Writings 5 (44): 620, 1916 – Fig. 2, 4, 6.

Lectotype. USA. Washington: Clallam Co., Sequim, *Grant* (BPI 701884, selected here).

= *Exidia villosa* Neuhoff, Pilze Mitteleuropas 2: 22, 1935. Neotype. Norway. Oslo: Bygdøy, Dronningberget, *Tilia cordata*, 3.V.2016 *Spirin 10012** (O, selected here).

= *Tremella albida* f. *betulae* P. Karst., Fungi Fenniae Exsiccati #317, 1866. Lectotype. Finland.

Etelä-Häme: Tammela, Mustiala, *Betula* sp., ‘December’, *Karsten* (Fungi Fenniae Exsiccati #317) (H, selected here).

= *Exidia albida* ssp. *tuberculata* P. Karst., Bidrag Känned. Finlands Nat. Folk 48: 452, 1889.

Lectotype. Finland. Etelä-Häme: Tammela, Taipamala, *T. cordata*, 23.IX.1888 *Karsten 2145* (H, selected here).

Basidiocarps persistent, semitranslucent, gelatinous, easily cut by a razor blade, adpressed, orbicular, later pulvinate to cerebriform, fusing together, up to 20 cm in widest dimension and 2 cm thick. Margin first villose, white, detaching, later adnate, compact, more or less concolorous with

hymenium. Hymenial surface pale to bright ochraceous, occasionally fading to almost white, indistinctly tuberculate or furrowed. Mineral inclusions often abundant, easily visible by a naked eye.

Hyphal structure monomitic; hyphae clamped (clamps often open), hyaline or brownish, often slightly thick-walled and 3–5 μm in diam. in deeper layers, more or less thin-walled and 2–4 μm in diam. in subhymenium, embedded in gelatinous matrix. Basidia four-celled, ellipsoid-ovoid, 11–16 \times 8.5–11 μm . Hyphidia branched, thin- to slightly thick-walled, hyaline to brownish, 0.5–2 μm in diam., spread among basidia and basidioles; epihymenial membrane absent. Cystidia-like cells abundant in vigorously growing basidiocarps, subfusiform or clavate, 15–25 \times 5–7 μm .

Basidiospores cylindrical, slightly or distinctly curved, (9.1–) 9.2–15.2 (–15.6) \times (3.1–) 3.2–4.8 (–5.0) μm ($n = 90/3$), $L = 11.41$, $W = 4.07$, $Q = 2.60$ –3.12.

Remarks. This species was first introduced as *Tremella albida* f. *betulae* (Karsten 1866). Later Karsten (1889) described it as *Exidia albida* ssp. *tuberculata*. Both of these taxa are typified here. Mussat (in Saccardo 1901) published a new combination, *E. tuberculata* (P. Karsten) Mussat, but he evidently did not accept it because this species epithet was treated as a synonym of Karsten's subspecies. Therefore, the species name *E. tuberculata* is considered invalid (Code Art. 36.1) (McNeill et al. 2012).

Lloyd (1916) described *E. candida* based on a single collection from Washington, USA. His description was amended by Burt (1921). The authentic specimen formally designated here as a lectotype is identical to three recent collections from the same region used in the present study.

Exidia villosa was described from East Prussia (Neuhoff 1935) and needs to be typified since original material is lacking. Fortunately, the protologue provides an excellent description of morphology and ecology of this taxon which is sufficient for selecting a neotype (indicated above). *Exidia candida* and *E. villosa* are morphologically indistinguishable, and DNA study confirms they are conspecific. Donk (1966) assumed that *Heterochaete europaea* Höhn. could be conspecific with

E. villosa. We studied the single original specimen of *H. europaea* (FH, selected here as a lectotype), and it certainly belongs to *Exidia thuretiana*.

Morphologically, *E. candida* is recognizable due to rather soft and often light-coloured basidiocarps, lack of epihymenial membrane and numerous mineral inclusions in hymenial layer. *Exidia thuretiana* is macroscopically similar but often totally white, and it can be distinguished by much larger basidiospores, $(12.2-13.8-18.2 (-18.7) \times (4.8-5.2-7.5 (-8.2) \mu\text{m}$ ($n = 60/2$), $L = 15.81$, $W = 6.27$, $Q = 2.35-2.74$. *Myxarium cinnamomescens* and *M. hyalinum* possess the same colours as those of *E. candida* but their basidiocarps are as a rule smaller and thinner; additionally, basidia of *Myxarium* spp. have a pronounced enucleate stalk.

In Europe, *E. candida* has more southern distribution than *E. candida* var. *cartilaginea*. It often occurs on just fallen branches of *Tilia cordata*, and can easily be detected on this substrate after strong winds. Some records were made on recently fallen lime logs. A few collections from other hosts exist, too. In North America, it has been found on *Corylus*.

Exidia candida var. *cartilaginea* (S. Lundell & Neuhoff) Spirin & V. Malysheva, stat. nov. – Fig. 2, 6.

≡ *Exidia cartilaginea* S. Lundell & Neuhoff, Pilze Mitteleuropas 2: 19, 1935. Lectotype. Sweden.

Uppland: Bondkyrka, Vårdsätra, *Betula* sp., 18.X.1932 Lundell (Fungi Exsiccati Suecici #263) (UPS F-010986, selected here).

MycoBank 821860

Basidiocarps persistent, semitranslucent, though gelatinous, often hardly cut by a razor blade, adpressed, orbicular, later pulvinate to somewhat cerebriform, fusing together, up to 5 cm in widest dimension and 1 cm thick. Margin first villose, white, detaching, later adnate, compact, paler than or more or less concolorous with hymenium. Hymenial surface pale ochraceous to bright

ochraceous-brown, then fading to almost white, indistinctly tuberculate to more or less clearly ridged. Mineral inclusions rare, seen as white grains under lens, or often absent.

Hyphal structure monomitic; hyphae clamped (clamps usually open), hyaline or brownish, often slightly thick-walled and 2.5–4.5 μm in diam. in deeper layers, more or less thin-walled and 2–3 μm in diam. in subhymenium, embedded in gelatinous matrix. Basidia four-celled, ellipsoid-ovoid, 11–17 \times 8–12 μm . Hyphidia branched, thin- to slightly thick-walled, hyaline to brownish, 1–2.5 μm in diam., densely arranged and glued together, producing firm layer on hymenial surface (epihymenial membrane). Cystidia-like cells abundant in vigorously growing basidiocarps, subulate or bottle-shaped, 15–25 \times 3–6 μm , sometimes producing simple or branched apical outgrowth. Basidiospores cylindrical, slightly to distinctly curved, (9.9–) 10.2–14.2 (–14.3) \times (3.8–) 4.0–5.1 (–5.2) μm ($n = 60/2$), $L = 11.90$, $W = 4.44$, $Q = 2.58$ –2.81.

Remarks. *Exidia cartilaginea* was described by Neuhoff (1935) based on several collections from Sweden. The type specimen is lost, and therefore we select another collection mentioned in the protologue (paratype) as a lectotype. As said above, *E. cartilaginea* should be considered merely a variety of *E. candida*.

One of two authentic specimens of *E. albida* subsp. *subrepanda* (Karsten 2142, on *Betula*, H) belongs to *E. candida* var. *cartilaginea*. This subspecies was raised to the species rank by Oudemans (1919: 398) but its identity remained obscure. In the protologue, Karsten (1891) stated *E. albida* subsp. *subrepanda* occurs on *Picea*. Therefore, we select another original specimen collected from spruce (Karsten 2143, H) as a lectotype of this taxon. This collection belongs to *Exidia saccharina* (Alb. & Schwein.) Fr., and thence we place *E. subrepanda* to the synonyms of the latter species.

Exidia candida var. *cartilaginea* differs from *E. candida* s. str. in having darker, sturdier basidiocarps with firm epihymenial membrane, and wider basidiospores. Most collections of *E. candida* var. *cartilaginea* are sterile or contain just a few mature basidiospores. Fertile specimens

are collected mostly in spring when the epihymenial membrane partly degrades and exposes basidia embedded in a dense, gelatinized layer of hyphidia.

This variety has a predominantly boreal distribution. It is often collected on dead but still attached branches of *Alnus* and *Betula* but found rarely also on other host species (*Acer*, *Salix*, *Tilia*). Here we confirm its presence in East Asia (Russian Far East) and the American North-West.

Myxarium cinnamomescens (Raitviir) Raitviir, Plants and Animals of the Far East: 113, 1971 – Fig. 3, 6.

≡ *Exidia cinnamomescens* Raitviir, Tartu Riikl. Ülikooli Toim. 136: 208, 1963. Holotype. Russia. Komi Rep.: Syktyvdinsky Dist., Graddor, *Populus tremula* (fallen log), 8.VIII.1957 Põldmaa (TAAM 5995, studied).

Basidiocarps persistent, gelatinous, first semitranslucent, pustulate, about 1 mm in diam., then fusing together, cerebriform or adpressed-orbicular, finally opaque, up to 8 mm in widest dimension and 1–3 mm thick. Margin detaching, later adnate, compact, more or less concolorous with hymenium. Hymenial surface hyaline with light yellowish tints, then almost white, indistinctly furrowed to more or less smooth, in herbarium specimens pale ochraceous or brownish. Mineral inclusions usually absent or, if present, very small, embedded in hymenial layer and thus seen only under lens.

Hyphal structure monomitic; hyphae clamped (clamps often open), hyaline, thin- or slightly thick-walled and 3–6 µm in diam. in deeper layers, at some septa swollen up to 9 µm in diam., thin-walled and 1–2.5 µm in diam. in subhymenium. Basidia four-celled, ellipsoid-ovoid to subglobose, 11–17.5 × 7–13 µm, with enucleate stalk up to 65 × 3.5 µm. Hyphidia branched, thin-walled, hyaline, 1–2 µm in diam., covering hymenial surface. Cystidia absent. Basidiospores cylindrical to broadly cylindrical, slightly to distinctly curved, (9.3–) 10.2–14.0 (–14.5) × (4.0–) 4.1–5.7 (–5.8) µm (n = 132/4), L = 11.96, W = 4.82, Q = 2.43–2.53.

Remarks. First described from Komi (European part of Russia) as *Exidia cinnamomeszens* (Raitviir 1963), the species was later moved to *Myxarium* (Raitviir 1971). Roberts (1998) suggested the synonymy with *Exidia nucleata* (= *Myxarium nucleatum*). However, as we show in this study, the older concept of *M. nucleatum* covers at least four species. The type of *M. cinnamomeszens* has been studied by us, and it is morphologically identical to two sequenced collections from Finland and Norway.

Macroscopically, *M. cinnamomeszens* is most similar to *M. populinum*, and these species can surely be separated only due to different size (especially width) of basidiospores. *Myxarium hyalinum* has more deeply coloured and larger basidiocarps that in the end of the growing season show abundant mineral inclusions, well visible through the almost translucent hymenial surface. In turn, *M. cinnamomeszens* never produces inclusions detectable by the naked eye, and its senescent basidiocarps become completely white and opaque. Moreover, basidiospores of *M. hyalinum* are on average wider than in *M. cinnamomeszens* although attention should be paid to the age of basidiocarps: younger specimens of *Myxarium* spp. tend to produce basidiospores somewhat narrower than in well developed fruitbodies.

Myxarium cinnamomeszens is distributed in the hemiboreal zone of Europe. As a rule, it inhabits still attached or just fallen branches of various deciduous trees. Its basidiocarps typically occur in ruptures of the bark, especially in those ones produced by stromatic pyrenomycetes. The other three gelatinous *Myxarium* species treated below usually grow on still corticated branches and logs. Here *M. cinnamomeszens* is reported from Finland, Norway and North-West Russia.

Myxarium hyalinum (Pers.) Donk, Persoonia 4: 233, 1966 – Fig. 3, 6.

≡ *Tremella hyalina* Pers., Mycol. Europea 1: 105, 1822. Neotype. Netherlands. Zuid-Holland: Leiden, *Acer* sp., 20.XI.1966 *Maas Geesteranus 15003** (H ex L) (selected here).

Basidiocarps persistent, semitranslucent, gelatinous, first adpressed-orbicular, often cerebriform, 2–5 mm in diam., then fusing together, up to 3 cm in widest dimension and 3 mm thick. Margin detaching, later adnate, compact, more or less concolorous with hymenium. Hymenial surface ochraceous to brownish, then fading to whitish (but staying translucent), indistinctly furrowed to more or less smooth, in herbarium specimens dark amber to brown. Mineral inclusions appearing only in coalescent basidiocarps, abundant, up to 1 mm in diam.

Hyphal structure monomitic; hyphae clamped (clamps often open), hyaline, thin- or slightly thick-walled and 2–4 μm in diam. in deeper layers, at some septa swollen up to 9 μm in diam., thin-walled and 1–2.5 μm in diam. in subhymenium. Basidia four-celled, ellipsoid-ovoid to subglobose, 11–18 \times 9–12 μm , with enucleate stalk up to 60 \times 3.5 μm . Hyphidia branched, thin-walled, hyaline, 1–2 μm in diam., covering hymenial surface. Basidiospores cylindrical to broadly cylindrical, slightly or distinctly curved, (9.0–) 9.2–15.7 (–16.7) \times (4.0–) 4.1–6.0 (–6.2) μm ($n = 90/3$), $L = 12.01$, $W = 5.01$, $Q = 2.38$ –2.45.

Remarks. The protologue of *T. hyalina* (Persoon 1822) is very concise and thus provides opportunities for different interpretations. Bourdot and Galzin (1909, 1928) were seemingly the first authors who adopted Persoon's species name for collections of *E. gemmata* (at that time a collective name for all gelatinous *Myxarium* species) lacking mineral inclusions. Neuhoﬀ (1935) accepted this opinion, albeit with some reservations. Following this line, Donk (1966) combined *T. hyalina* to *Myxarium* in order to replace *M. nucleatum* with an older name. Reid (1970) argued that Persoon's description is so inexact that it could hardly be applied to any specific species. Roberts (1998) disagreed with both Donk's synonymy and his elimination of *M. nucleatum* from *Exidia*. At the same time, Mycobank database (www.mycobank.org) lists *M. hyalinum* as having priority over *M. nucleatum*.

We agree that Persoon's description is open to various interpretations, and therefore our solution for this problem is merely pragmatical – to avoid as many name changes as possible. We studied and

sequenced four morphologically identical collections including one identified by Donk as *M. hyalinum* (Maas Geesteranus 15003). Three of them lack mineral stuff while the specimen labelled by Donk represents senescent, coalescent basidiocarps with numerous, well-visible inclusions. Since the Central European species is dealt with, we decided to name it *Myxarium hyalinum*. Our solution at least partly corresponds to the idea of the species developed by Bourdot and Galzin and their followers. A neotype of *T. hyalina* from the Netherlands is designated here. Normally developed basidiocarps of *M. hyalinum* differ from other *Myxarium* species in being more robust and having deeper, ochraceous or brownish colours. Basidiospores and basidia of *M. hyalinum* are on average larger than in *M. nucleatum* and *M. populinum*. Differences of *M. hyalinum* from *M. cinnamomesceus* are discussed under remarks to the latter species. *Myxarium hyalinum* is a southern species distributed mainly in the nemoral zone. We studied specimens from Denmark, Germany, the Netherlands, Norway and Ukraine.

Myxarium nucleatum Wallr., Flora Cryptogam. Germaniae 2: 260, 1833 – Fig. 3, 5, 6.

Neotype. Norway. Oslo: Bygdøy, Dronningberget, *Tilia cordata*, 3.V.2016 *Spirin 10013** (O, selected here).

Basidiocarps persistent, translucent, gelatinous, easily cut by a razor blade, adpressed-orbicular, 1–5 mm in diam., often in gregarious groups, partly fusing together, 0.1–0.5 mm thick, drying to almost invisible vernicose films. Margin adnate, compact, more or less concolorous with hymenium. Hymenial surface whitish or faintly yellowish to pale ochraceous, more or less smooth. Mineral inclusions always present, abundant, 0.4–0.8 mm in diam.

Hyphal structure monomitic; hyphae clamped (clamps often open), hyaline, thin- or slightly thick-walled and 2–3.5 µm in diam. in deeper layers, at some septa swollen up to 7 µm in diam., thin-walled and 1–2 µm in diam. in subhymenium. Basidia four-celled, ellipsoid-ovoid to subglobose, 10.5–13 × 8–10.5 µm, with enucleate stalk up to 30 × 2.5 µm. Hyphidia branched, thin-walled,

hyaline, 1–2 µm in diam., covering hymenial surface. Basidiospores cylindrical, slightly or distinctly curved, (7.9–) 8.5–13.8 (–14.0) × (3.2–) 3.5–5.2 (–5.4) µm (n = 130/4), L = 10.88, W = 4.26, Q = 2.38–2.70.

Remarks. Wallroth (1833) described *Myxarium* as monotypic, with *M. nucleatum* as the only accepted species. His description contains a reference to the North American *Tremella nucleata* Schwein. (as ‘*Nematelia nucleata* Fr.’). However, as Donk (1966) correctly noted, Wallroth’s *M. nucleatum* is not a simple recombination of Schweinitz’s species to the new genus but a new species name referring to a different species. No type exists for *M. nucleatum*, and therefore we designate a neotype here. In our opinion, pale-coloured, adpressed, coalescent basidiocarps with numerous, distinctive mineral inclusions are rather clear indications of what species was dealt with in the protologue.

Basidiocarps of *M. nucleatum* are hardly visible in dry condition and detectable mainly due to a presense of white mineral grains. This feature makes *M. nucleatum* different from the microscopically similar *M. populinum*, as well as from *M. cinnamomescens*. The latter species also possesses larger basidia and basidiospores. Senescent basidiocarps of *M. hyalinum* may also bear mineral inclusions but the latter species has more deeply coloured basidiocarps, well visible in dry condition, and bigger basidiospores.

Tremella gemmata Lév. was described from Crimea (Demidov 1842), and it is often considered a synonym of *M. nucleatum*. The protologue points towards the latter species as accepted here.

However, this synonymy should be re-confirmed. *Exidia alboglobosa* Lloyd was also listed among synonyms of *M. nucleatum* (Reid 1970) but its identity should be re-established. White globose basidiocarps are untypical for *M. nucleatum* s.str., and they may well represent the senescent, discoloured specimen of *M. cinnamomescens* or *M. hyalinum*. This problem, however, can be solved with a newly collected and sequenced material from France.

Myxarium nucleatum seems to be widely distributed in temperate zone although its distribution area stretches up to the Arctic Circle, with the northernmost locality in Kuusamo (Finland). The favorite host of *M. nucleatum* in North Europe is *Tilia cordata*. Here the species is reported from Denmark, Finland, Norway, North-West Russia, and Sweden.

Myxarium populinum (P. Karst.) Spirin & V. Malysheva, comb. nov. – Fig. 3, 6.

≡ *Exidia albida* var. *populina* P. Karst., Medd. Soc. Fauna Flora Fennica 18: 73, 1891. Lectotype. Finland. Etelä-Häme: Tammela, Mustiala, *Populus tremula*, XI.1890 *Karsten 1812* (H, selected here).

MycoBank 821861

Basidiocarps persistent, gelatinous, first semitranslucent, adpressed-orbicular, about 3 mm in diam., then fusing together, up to 15 mm in widest dimension and 0.5–1 mm thick. Margin detaching, compact, more or less concolorous with hymenium. Hymenial surface whitish, sometimes with light yellowish tints, indistinctly furrowed to more or less smooth, in herbarium specimens pale ochraceous or brownish. Mineral inclusions usually absent or, if present, very small, embedded in hymenial layer and thus seen only under lens.

Hyphal structure monomitic; hyphae clamped (clamps often open), hyaline, thin- or slightly thick-walled and 2.5–5.5 µm in diam. in deeper layers, at some septa swollen up to 8 µm in diam., thin-walled and 1–2.5 µm in diam. in subhymenium. Basidia four-celled, ellipsoid-ovoid to subglobose, 10.5–13 × 9–12 µm, with enucleate stalk up to 50 × 2.5 µm. Hyphidia branched, thin-walled, hyaline, 1–2 µm in diam., covering hymenial surface. Basidiospores cylindrical to narrowly cylindrical, slightly or distinctly curved, (8.7–) 9.0–13.8 (–16.2) × (3.0–) 3.1–4.9 (–5.0) µm (n = 150/5), L = 11.08, W = 3.94, Q = 2.72–2.95.

Remarks. This species was initially described as *E. albida* var. *populina* (Karsten 1891). Oudemans (1920: 94) raised this variety to the species rank. Karsten's single original collection is indicated

here as a lectotype. It is morphologically identical to a number of specimens collected in North Europe, of which three have been sequenced. From macroscopically similar *M. cinnamomescens* and *M. hyalinum*, this species differs in having considerably narrower basidiospores. *Myxarium nucleatum* possesses basidiospores of almost the same size as *M. populinum* but its basidiocarps look different.

Myxarium populinum seems to be rather common in North Europe. All but one collections studied by us came from aspen.

Specimens examined (sequenced specimens are marked by asterisk)

Exidia candida. Canada. British Columbia: Vancouver, deciduous tree, 15.IV.1968 *Bandoni** (LE 38198). Finland. Uusimaa: Helsinki, Käpylä, *Tilia cordata*, 30.IV.2015 *Miettinen* 19135 (H). Etelä-Häme: Tammela, Mustiala (H, lectotype of *Tremella albida* f. *betulae*, see above), Taipamala (H, lectotype of *T. albida* subsp. *tuberculata*, see above). Norway. Oslo: Bygdøy (O, neotype of *Exidia villosa*, see above). Vest-Agder: Flekkefjord, Hidra, *T. cordata*, 16.IV.1973 *Gulden* 439/73 (O F-179584). Russia. Khabarovsk Reg.: Solnechnyi Dist., Igdomi, *Betula platyphylla*, 7.VIII.2011 *Spirin* 3921* (H). Leningrad Reg.: Lodeinoe Pole Dist., Nizhnesvirsky Nat. Res., *Alnus incana*, 18.IX.1996 *Zmitrovich** (LE 201847). Mordovia: Temnikov Dist., Mordovsky Nat. Res., *T. cordata*, 2.X.2014 *Bolshakov** (LE 313211). Nizhny Novgorod Reg.: Lukoyanov Dist., Razino, *T. cordata*, 6.VIII.2013 *Spirin* 5999 (H), 7–8.VIII.2014 *Spirin* 7212*, 7263 (H), 19–21.VIII.2015 *Spirin* 9595, 9761 (H), Srednii, *T. cordata*, 18.VIII.2006 *Spirin* 2600 (H). USA. Washington: Clallam Co. (lectotype of *E. candida*, see above); Pend Oreille Co., Lime Creek, *Corylus cornuta*, 15.X.2014 *Spirin* 8584*, 8588* (H).

Exidia candida var. *cartilaginea*. Finland. Etelä-Häme: Tammela, Mustiala, *Betula* sp., 18.XII.1890 *Karsten* 2142 (H, as *E. albida* subsp. *subrepanda*). Norway. Baerum: Kjaglidalen, *A. incana*, 4.V.2016 *Spirin* 10021* (O); Lysakerelven, *Acer pseudoplatanus*, 16.X.1964 *Torkelsen* 159/4 (O

F160479). Hordaland: Kvinnherad, Stonganeset, *T. cordata*, 27.XII.1984 *Flatabø** (O F-160269).
 Russia. Khabarovsk Reg.: Solnechnyi Dist., Igdomi, *Alnus hirsuta*, 2.IX.2016 *Spirin 10762** (H),
 3.IX.2016 *Spirin 10906* (H), *Alnus (Duschekia) fruticosa*, 4.IX.2016 *Spirin 10954** (H), Suluk-
 Makit, *Salix* sp., 17.VIII.2011 *Spirin 4159* (H); Verkhnebureinskii Dist., Hegdy, *A. fruticosa*,
 22.VIII.2014 *Spirin 7833* (H). Krasnoyarsk Reg.: Ermakovskoe Dist., Sayano-Shushensky Nature
 Reserve, *Alnus* sp., 17.VIII.2015 *Malysheva** (LE 312774). Leningrad Reg.: Podporozhie Dist.,
 Vazhinka, *Betula pubescens*, 21.V.2016 *Spirin 10105** (H). Republic of Altai: Ulagan Dist., Altai
 Nature Reserve, 18.VIII.2008 *Malysheva** (LE 231797). Sweden. Uppland: Bondkyrka, Vårdsätra,
 (UPS F-010986, lectotype of *E. cartilaginea*, see above). USA. Washington: Pend Oreille Co.,
 Sullivan Creek, *Alnus rubra*, 13.X.2014 *Spirin 8450** (H).

Exidia saccharina. Finland. Etelä-Häme: Tammela, Mustiala, *Picea abies*, 15.XI.1890 *Karsten*
2143 (H, lectotype of *E. subrepanda*).

Exidia thuretiana. Bosnia and Herzegovina. Jaize, *Fagus sylvatica*, 14.IV.1901 *Höhnelt* (FH –
 lectotype of *Heterochaete europaea*). Finland. Uusimaa: Helsinki, Viikki, *Ribes nigrum*,
 28.IV.2016 *Spirin 9999** (H). Norway. Akershus: Baerum, Kjaglidalen, *Ulmus glabra*, 16.IX.2016
Spirin 11128 (O). Telemark: Nome, Mörkvasslia Nat. Res., *U. glabra*, 25.X.2016 *Spirin 11185**
 (O). Russia. Leningrad Reg.: Podporozhie Dist., Grishino, *Ribes* sp., 26.V.2017 *Spirin 11207* (H).
Myxarium cinnamomescens. Finland. Etelä-Häme: Kangasala, Vihtinen, *Padus avium*, 7.IX.2008
*Söderholm 3990** (H). Norway. Aust-Agder: Arendal, Tromøy, *T. cordata*, 25.XII.1972 *Pedersen*
 (O F160483). Akershus: Ås, Ved Høyskolen, *Populus tremula*, 10.XII.1972 *Hansen & Gulden*
*595/72** (O F160494); Bærum, Eikeli, *Betula* sp., 26.X.1971 *Schumacher* (O F160499),
 Sandvikselva, garden shrub, 22.V.1972 *Schumacher* (O F160498). Oslo: Bygdøy, *T. cordata*,
 9.XI.1969 *Eftenstøl* (O F160490). Buskerud: Hole, Vik, *Betula* sp., 13.IX.1965 *Torkelsen* (O
 F160489), 30.IX.1967 *Torkelsen* (O F160484). Hedmark: Hamar, Domkirkeodden, *Betula*
pubescens, 19.IX.1965 *Torkelsen 135/5* (O F160491). Hordaland: Ulvik, Osa, *Betula* sp.,

3.VII.1951 *Stordal* (O F160502). Oppland: Etnedal, Bruflat i Valdres, *P. tremula*, *Blytt* (O F160503). Russia. Komi Rep. (holotype, see above).

Myxarium hyalinum. Denmark. Sjælland: Slagelse, Borreby, *T. cordata*, 14.IV.2012 *Læssøe TL2012-443455** (C, dupl. H). Germany. Bremen: Botanical Garden, deciduous tree, 28.VI.2012 *Miettinen 15436.2** (H). Nordrhein-Westfalen: Bielefeld, Ochsenburg, *Fagus sylvatica*, 25.XI.2015 *Miettinen 19855** (H). Netherlands. Zuid-Holland: Leiden (neotype, see above). Norway. Vest-Agder: Flekkefjord, Hidra, *Quercus robur*, 16.IV.1973 *Gulden 431/73* (O F160501). Akershus: Bærum, Evje, *Fraxinus excelsior*, 27.VII.1952 *Stordal* (O F160485). Ukraine. Kharkov Reg.: Kharkov, Forest Park, *Acer platanoides*, *Q. robur*, 25.IX.2013 *Akulov* (CWU).

Myxarium nucleatum. Denmark. Jylland: Horsens, Hansted skov, *F. sylvatica*, 19.IX.1972 *Torkelsen* (O F160480). Finland. Etelä-Häme: Lahti, Ritämäki, *T. cordata*, 26.X.1998 *Haikonen 19456* (H). Koillismaa: Kuusamo, Liikasenvaara, *Ribes* sp., 23.VIII.1978 *Ryvarden* (O F160841). Norway. Vestfold: Tønsberg, Slagentangen, *T. cordata*, 24.I.1993 *Marstad* (O F160504), 25.I.1993 *Marstad 42-93** (O F165125). Akershus: Baerum, Lysakerelven, *A. pseudoplatanus*, 16.X.1964 *Torkelsen 158/4* (O F160478). Oslo: Bygdøy, Dronningberget (neotype, see above). Aust-Agder: Grimstad, Reddalskollen, *T. cordata*, 11.IV.2005 *Fonneland 05-01* (O F157942); Bygland, Fånefjell, *T. cordata*, 1.X.2006 *Fonneland 06-144* (O F282994). Møre og Romsdal: Averøy, Mellom Bodalen og Hoset, *Corylus avellana*, 4.VIII.1974 *Torkelsen 779/74* (O F160477). Russia. Leningrad Reg.: Tosno Dist., *T. cordata*, 2.X.1997 *Zmitrovich** (LE 206820). Sweden. Uppland: Bondkyrka, Hällby, *P. tremula*, 15.X.1944 *Nannfeldt 7532* (O F160508).

Myxarium populinum. Finland. Uusimaa: Nurmijärvi, Kirkonkylä, *P. tremula*, 21.XI.1982 *Askola 1185** (H). Etelä-Häme: Nastola, Villähde, *P. tremula*, 13.XI.2005 *Haikonen 24623** (H); Tammela, Mustiala (lectotype of *Exidia albida* var. *populina*, see above). Norway. Vestfold: Larvik, Jordstøyp i Kvelde, *P. tremula*, 6.IX.1995 *Andersen** (O F90775). Akershus: Nannestad, Hurdalssjøen, *P. tremula*, 28.IX.1969 *Gulden* (O F160506). Buskerud: Kongsberg, Fulsebakkåsen i

Jondalen, *P. tremula*, 21.VIII.1999 *Winter* (O F63156); Nes, Eidal i Rukkedalen, *Salix caprea*, 2.X.1965 *Torkelsen 151/5* (O F160500). Hedmark: Ringsaker, Veldre, *P. tremula*, 19.IX.1965 *Torkelsen 136/5* (O F160507).

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Figures

Fig. 1. Phylogenetic relationships of *Exidia* and *Myxarium* species. Collections are indicated with the voucher number: a – the best tree from the ML analyses of the combined dataset

nrITS+nrLSU; b – the best tree from the ML analyses of the *tef1* dataset. Support values (ML/BA) are given on the branches. Sequences of *Exidia candida* var. *cartilaginea* are marked by asterisk.

Fig. 2. Basidiocarps: a, b – *Exidia candida* (Spirin 9761, LE 313211); c, d – *E. candida* var. *cartilaginea* (Spirin 10954, LE 231797); e, f – *E. thuretiana* (Spirin 9999). Scale bar = 5 mm

Fig. 3. Basidiocarps: a – *Myxarium populinum* (Haikonen 24623); b – *M. cinnamomescens* (Hansen 595/72); c – *M. nucleatum* (LE 206820) in fresh condition; d – *M. nucleatum* (LE 206820) in dry condition; e, f – *M. hyalinum* (Miettinen 19855, Miettinen 15436.2). Scale bar = 5 mm.

Arrows point to white mineral grains.

Fig. 4. Microscopic structures of *Exidia candida* (Spirin 10012). Scale bar = 10 µm

Fig. 5. Microscopic structures of *Myxarium nucleatum* (Spirin 10013). Scale bar = 10 µm

Fig. 6. Basidiospores of *Exidia* and *Myxarium* spp.: a – *Exidia candida* var. *cartilaginea* (Spirin 10021), b – *E. candida* (Spirin 10012), c – *Myxarium hyalinum* (Maas Geesteranus 15003), d – *M. cinnamomescens* (Hansen 595/72), e – *M. nucleatum* (Spirin 10013), f – *M. populinum* (Karsten 1812). Scale bar = 5 µm

Table 1

Table 1. Data for ITS and LSU sequences used in the phylogenetic analyses

Species	Collector / herbarium number	Origin (ISO code)	Host	LSU GenBank #	ITS GenBank #	<i>tef 1</i> GenBank #	Source
<i>Exidia candida</i> *	O F160269	NO	<i>Tilia</i>	KY801897	KY801872	KY801922	this study
<i>E. candida</i> *	Spirin 8450 (H)	US-WA	<i>Alnus</i>	KY801900	KY801875	KY801925	this study
<i>E. candida</i> *	Spirin 10954 (H)	RU- KHA	<i>Alnus</i>	KY801901	KY801876	KY801926	this study
<i>E. candida</i>	LE 38198	CA-BC	-	KY801896	KY801871	KY801921	this study
<i>E. candida</i>	Spirin 8588 (H)	US-WA	<i>Corylus</i>	KY801895	KY801870	KY801920	this study
<i>E. candida</i> *	LE 312774	RU- KYA	<i>Alnus</i>	KY801902	KY801877	-	this study
<i>E. candida</i>	Spirin 3921 (H)	RU- KHA	<i>Betula</i>	KY801892	KY801867	KY801918	this study
<i>E. candida</i> *	LE 231797	RU-AL	-	KY801904	-	-	this study
<i>E. candida</i> *	Spirin 10021 (O)	NO	<i>Alnus</i>	KY801899	KY801874	KY801924	this study
<i>E. candida</i>	LE 313211	RU-MO	<i>Tilia</i>	KY801893	KY801868	KY801919	this study
<i>E. candida</i>	LE 201847	RU- LEN	<i>Alnus</i>	KY801903	-	-	this study
<i>E. candida</i> *	Spirin 10105 (H)	RU- LEN	<i>Betula</i>	KY801898	KY801873	KY801923	this study
<i>E. candida</i>	Spirin 10012 (O)	NO	<i>Tilia</i>	KY801890	KY801865	KY801916	this study
<i>E. candida</i>	Spirin 7212 (H)	RU-NIZ	<i>Tilia</i>	KY801891	KY801866	KY801917	this study
<i>Exidia japonica</i>	TAAM 42689	RU-PRI	-	AF291320	AF291274	-	Weiss & Oberwinkler (2001)
<i>Exidia uvapassa</i>	AFTOL-ID 461		-	AY645056	DQ241776	-	GenBank
<i>E. uvapassa</i>	TUFC 34007	JP	-	AB871843	AB871863	-	GenBank
<i>E. truncata</i>	MW 365	DE	-	AF291325	AF291279	-	Weiss & Oberwinkler (2001)

<i>E. recisa</i>	MW 315	DE	-	AF291322	AF291276	-	Weiss & Oberwinkler (2001)
<i>E. thuretiana</i>	Spirin 9999 (H)	FI	<i>Ribes</i>	KY801905	KY801878	KY801927	this study
<i>E. thuretiana</i>	GEL 5242	DE	-	AF291278	AF291278	-	GenBank
<i>E. thuretiana</i>	Spirin 11185 (O)	NO	<i>Ulmus</i>	KY801914	KY801889	-	this study
<i>E. thuretiana</i>	MW 373	DE	-	AF291324	AF291278	-	Weiss & Oberwinkler (2001)
<i>Exidia saccharina</i>	RoKi 88	DE	-	AF291323	AF291277	-	Weiss & Oberwinkler (2001)
<i>Exidia pithya</i>	MW 313	DE	-	AF291321	AF291275	-	Weiss & Oberwinkler (2001)
<i>Exidia glandulosa</i>	MW 355	DE	-	AF291319	AF291273	-	Weiss & Oberwinkler (2001)
<i>Myxarium cinnamomescens</i>	O F160494	NO	<i>Populus</i>	KY801909	KY801882	-	this study
<i>M. cinnamomescens</i>	Söderholm 3990 (H)	FI	<i>Padus</i>	KY801912	KY801886	-	this study
<i>Myxarium populinum</i>	Haikonen 24623 (H)	FI	<i>Populus</i>	KY801910	KY801883	-	this study
<i>M. populinum</i>	Askola 1185 (H)	FI	<i>Populus</i>	KY801911	KY801885	-	this study
<i>M. populinum</i>	O F90775	NO	<i>Populus</i>	-	KY801888	-	this study
<i>Myxarium subhyalinum</i>	MW 527	DE	-	AF291352	-	-	Weiss & Oberwinkler (2001)
<i>Myxarium granulum</i>	USJ 54532	CR	-	AF291348	-	-	Weiss & Oberwinkler (2001)
<i>Myxarium grilletii</i>	RoKi 218	DE	-	AF291349	-	-	Weiss & Oberwinkler (2001)
<i>Myxarium nucleatum</i>	Spirin 10013 (O)	NO	<i>Tilia</i>	KY801906	KY801879	-	this study
<i>M. nucleatum</i>	LE 206820	RU-	<i>Tilia</i>	KY801894	KY801869	-	this study

		LEN					
<i>M. nucleatum</i>	O F165125 (H)	NO	<i>Tilia</i>	-	KY801884	-	this study
<i>Myxarium</i> <i>mesonuclearum</i>	USJ 55354	CR	-	AF291350	-	-	Weiss & Oberwinkler (2001)
<i>M. hyalinum</i>	Geesteranus 15003 (H)	NL	<i>Acer</i>	KY801908	KY801881	-	this study
<i>M. hyalinum</i>	Miettinen 15436.2 (H)	DE	-	KY801913	KY801887	-	this study
<i>M. hyalinum</i>	TL2012-443455 (C)	DK	<i>Tilia</i>	KY801907	KY801880	KY801928	this study
<i>M. hyalinum</i> **	ZP TRE2M	PT	-	AF291351	-	-	Weiss & Oberwinkler (2001)
<i>M. hyalinum</i>	Miettinen 19855 (H)	DE	<i>Fagus</i>	KY801915	-		this study
<i>M. nucleatum</i> s.l.	KW 3885	US-CA	-	AY509554	AY509554	-	Wells et al. (2004)

* *Exidia candida* var. *cartilaginea*

** Published as *M. nucleatum* (Weiss & Oberwinkler 2001)